Bayesian Hierarchical MPT Models Application and Practice with TreeBUGS

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TreeBUGS: Bayesian Hierarchical MPT Modeling

- The R package TreeBUGS
- Basic modeling
 - Model fitting
 - Convergence
 - Plots
 - Model fit
- 3) Advanced modeling
 - Within-subject comparisons
 - Between-subject comparisons
 - Covariates
- Sensitivity/robustness analysis
 - Priors
 - Predictive distributions
 - Simulation

Software for Hierarchical MPT Models

Software for Hierarchical MPTs

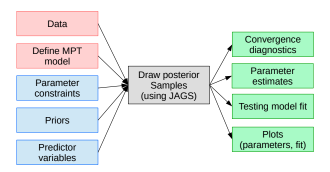
- Implementation of MCMC sampling in R/C/Fortran (Klauer 2010)
- General-purpose software: WinBUGS/JAGS/Stan (Matzke et al. 2015)
- Requires re-implementation of summaries, statistics, plots

TreeBUGS: A user-friendly R package (Heck, Arnold, and Arnold 2018)

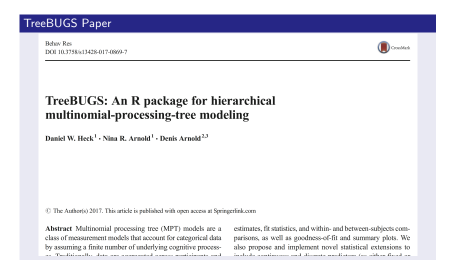
- Easy-to-use, open source, free
- Fitting and testing MPT models
 - Posterior sampling, summary statistics, and plots
 - Data generation, robustness simulations
 - Change priors, add predictors, etc.
 - Current limitation: Crossed random effects (persons + items)

Functionality of TreeBUGS

- Input: R objects or text/csv files (minimal R knowledge required)
- Priors and other details can be changed in R
- TreeBUGS translates the model to JAGS (Plummer, 2003) to draw posterior samples
- Functions for post-processing, summaries and plots



Reference



Basic Modeling

(corresponding R script: 07-ApplicationIII-TreeBUGS.R)

MPT Model Specification

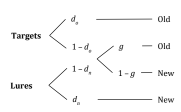
- MPT structure is defined in an EQN model file
 - Simple: copy from multiTree :-)
- Difference: the symbol # allows to add comments

Two-high threshold model

```
(file: 2htm.eqn)
```

```
# Targets
target hit do
target hit (1-do)*g
target miss (1-do)* (1-g)

# Lures
lure cr dn
lure fa (1-dn)*g
lure cr (1-dn)*(1-g)
```



Data Structure

- Data: Response frequencies in wide format
 - One line per person
 - One category per column
 - Column names must be identical to the EQN categories!
- Either supplied in .csv-file or as data.frame / matrix in R

Example: 2htm.csv

```
# set working directory:
# setwd("D:/R/MPT-workshop/")
frequencies <- read.csv("2htm.csv")
head(frequencies, 5)</pre>
```

```
## cr fa hit miss
## 1 26 24 32 18
## 2 46 4 45 5
## 3 36 14 38 12
## 4 46 4 38 12
```

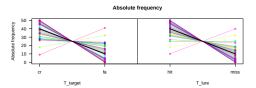
Heterogeneity

- Load TreeBUGS and plot heterogeneity
- If response frequencies are homogeneous, standard (fixed-effects) MPT models are statistically more efficient

```
library("TreeBUGS")
plotFreq(frequencies, eqnfile = "2htm.eqn")
```



```
plotFreq(frequencies, boxplot = FALSE, eqnfile = "2htm.eqn")
```



Fitting MPT Models

- Fitting an MPT model in TreeBUGS
 - Model: Text file in EQN syntax (with model equations)
 - Data: .csvfile
 - Constraints: text file with equality contraints

Fitting an MPT Model in R

- Alternative: define everything directly in R
 - Model: Text string (character in apostrophes)
 - Data: Matrix or data frame

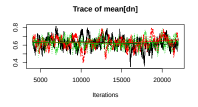
■ Constraints: A list

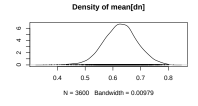
Equality constraints

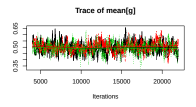
```
# (A) use a general model file and constrain parameters:
fit <- traitMPT(eqnfile = htm,</pre>
                data = frequencies,
                restrictions = list("dn=do", "g=.50"))
# (B) hard-coding of constraints in the EQN file:
htm constr <- "
target hit d
target hit (1-d)*.50
target miss (1-d)*.50
lure cr d
lure fa (1-d)*.50
lure cr (1-d)*.50
fit <- traitMPT(eqnfile = htm_constr,</pre>
                data = frequencies)
```

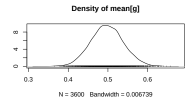
- Convergence check
 - Posterior/ MCMC samples should look unsystematic (like a hairy caterpillar)
 - For more options, see: ?plot.traitMPT

plot(fit, parameter = "mean", type = "default")

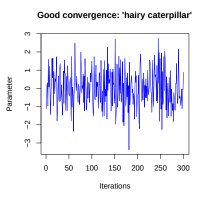




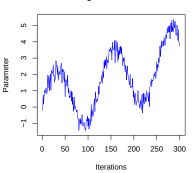




■ Interpreting MCMC plots



Bad convergence: slow movement

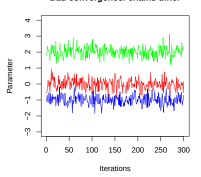


- Gelman-Rubin statistic (\hat{R})
 - Also known as: "potential scale reduction factor" or "R hat"
 - Similar to ANOVA: Compares between-chain and within-chain variances (large differences between these variances indicate nonconvergence)
 - Statistic should be close to 1 (standard criterion: $\hat{R} < 1.05$)

Good convergence: all chains similar

Paramore Company of the Paramo

Bad convergence: chains differ



summarv(fit)

- Gelman-Rubin statistic (\hat{R})
 - Columns Rhat and R_95% in the summary output

```
## Call:
    ## traitMPT(eqnfile = htm, data = frequencies, restrictions = list("dn=do"),
    ##
          ppp = 1000)
    ##
    ## Group-level medians of MPT parameters (probability scale):
    ##
               Mean
                      SD 2.5% 50% 97.5% Time-series SE n.eff Rhat R 95%
    ## mean dn 0.627 0.061 0.504 0.628 0.745
                                                  0.004
                                                          239 1.018 1.053
    ## mean_g 0.498 0.041 0.419 0.498 0.581
                                                  0.002 449 1.021 1.073
    ##
    ## Mean/Median of latent-trait values (probit-scale) across individuals:
    ##
                    Mean
                            SD
                                 2.5% 50% 97.5% Time-series SE n.eff Rhat
    ## latent mu dn 0.328 0.163 0.011 0.325 0.658
                                                          0.011
                                                                 237 1.017
    ## latent mu g -0.005 0.104 -0.205 -0.006 0.204
                                                         0.005 450 1.021
    ##
                  R 95%
    ## latent mu dn 1.053
    ## latent mu g 1.073
    ##
    ## Standard deviation of latent-trait values (probit scale) across individuals:
                              SD 2.5% 50% 97.5% Time-series SE n.eff Rhat
    ##
                      Mean
    ## latent sigma dn 1.120 0.146 0.872 1.107 1.442 0.003 2534 1.005
    ## latent sigma g 0.625 0.096 0.462 0.616 0.836 0.002 2434 1.002
    ##
                     R 95%
    ## latent_sigma_dn 1.019
## latent_sigma_g 1.008
```

Options for MCMC Sampling

If the model has not converged, it must be fitted with more conservative settings:

```
fit <- traitMPT(
   eqnfile = htm, data = frequencies,
   restrictions = list("dn=do"),

n.adapt = 5000, # longer adaption of JAGS increases efficiency of sampling
   n.burnin = 5000, # longer burnin avoids issues due to bad starting values
   n.iter = 30000, # drawing more MCMC samples leads to higher precision
   n.thin = 10, # ommitting every 10th sample reduces memory load
   n.chains = 4) # more MCMC chains increase precision</pre>
```

```
## MCMC sampling started at 2019-04-30 09:36:44
## Calling 4 simulations using the parallel method...
## Following the progress of chain 1 (the program will wait for all
## chains to finish before continuing):
## Welcome to JAGS 4.3.0 on Tue Apr 30 09:36:48 2019
## JAGS is free software and comes with ABSOLUTELY NO WARRANTY
## Loading module: basemod: ok
## Loading module: bugs: ok
## . Loading module: dic: ok
## . Loading module: glm: ok
## . Reading data file data.txt
## . Compiling data graph
Daniel ##Heck Resolving undeclared variables
```

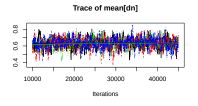
Extend MCMC Sampling

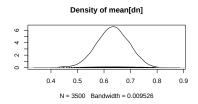
- If the MCMC samples are OK but higher precision is needed:
 - Extend sampling and add new MCMC samples to the fitted JAGS object

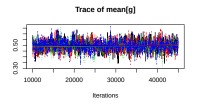
```
## Calling 4 simulations using the parallel method...
    ## Following the progress of chain 1 (the program will wait for all
    ## chains to finish before continuing):
    ## Welcome to JAGS 4.3.0 on Tue Apr 30 09:37:28 2019
    ## JAGS is free software and comes with ABSOLUTELY NO WARRANTY
    ## Loading module: basemod: ok
    ## Loading module: bugs: ok
    ## . Loading module: dic: ok
    ## . Loading module: glm: ok
    ## . . Reading data file data.txt
    ## . Compiling data graph
    ##
         Resolving undeclared variables
    ##
          Allocating nodes
    ##
          Initializing
    ##
          Reading data back into data table
    ## Compiling model graph
    ##
          Resolving undeclared variables
    ##
          Allocating nodes
    ## Graph information:
    ##
          Observed stochastic nodes: 100
## Unobserved stochastic nodes: 55
```

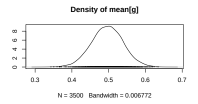
Convergence for Second Fit

■ Check convergence again:









Parameter Estimates

- Summary statistics for posterior distribution:
 - Posterior mean and median (50% quantile)
 - Posterior standard deviation (SD, similar to standard error)
 - Bayesian credibility interval (2.5% and 97.5% quantiles)

```
summary(fit2)
```

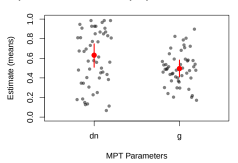
```
## Call:
    ## [[1]]
    ## traitMPT(eqnfile = htm, data = frequencies, restrictions = list("dn=do"),
           n.iter = 30000, n.adapt = 5000, n.burnin = 5000, n.thin = 10,
    ##
           n.chains = 4)
    ##
    ##
    ## [[2]]
    ## extendMPT(fittedModel = fit. n.iter = 10000, n.adapt = 2000,
    ##
           n.burnin = 0)
    ##
    ##
    ## Group-level medians of MPT parameters (probability scale):
                       SD 2.5% 50% 97.5% Time-series SE n.eff Rhat R_95%
    ##
                Mean
    ## mean_dn 0.633 0.061 0.512 0.633 0.748 0.002 675 1.002 1.004
    ## mean g 0.496 0.043 0.410 0.497 0.582
                                                 0.001 1288 1.003 1.009
    ##
    ## Mean/Median of latent-trait values (probit-scale) across individuals:
    ##
                             SD 2.5% 50% 97.5% Time-series SE n.eff Rhat
                     Mean
    ## latent mu dn 0.343 0.163 0.029 0.341 0.668 0.006 672 1.002
    ## latent mu g -0.010 0.110 -0.227 -0.008 0.207 0.003 1288 1.003
    ##
                   R 95%
    ## latent mu dn 1.005
    ## latent_mu_g 1.009
Daniel ## Standard deviation of latent-trait values (probit scale) across individuals:
```

Plot Parameter Estimates

- Estimates for group-level parameters
 - lacktriangle Overall mean μ : Posterior mean and Bayesian credibility interval
 - Individual parameters θ_i : Posterior mean

plotParam(fit)

roup-level means + 95% CI (red) and individual means

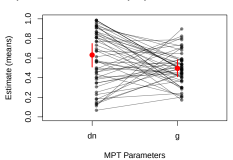


Plot Parameter Estimates

- Plot parameter profiles
 - E.g., assess test-retest reliability of a parameter (Michalkiewicz & Erdfelder, 2016)
- For more options, see: ?plotParam

```
plotParam(fit, addLines = TRUE, select = c("dn", "g"))
```

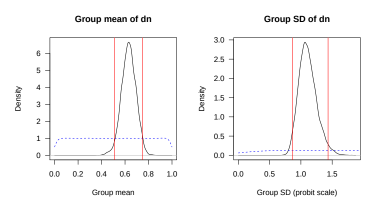
roup-level means + 95% CI (red) and individual means



Compare Prior and Posterior

How much did we learn about the parameters?

- Graphical assessment: Plot prior (blue) and posterior (black) densities plotPriorPost(fit)
- ## Press <Enter> to show the next plot.

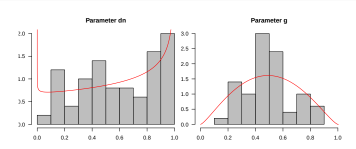


Group-Level Distribution

Distribution of individual estimates

- Histogram: Distribution of θ estimates (posterior mean per person)
- Red density: Estimated group-level distribution

plotDistribution(fit) # graphical test



Model Fit: Predicted vs. Observed Data

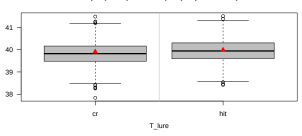
Graphical test of model fit

- Plot means of observed frequencies
- Compare against posterior-predicted frequencies (boxplots)

```
colMeans(frequencies) # observed group means that are tested

## cr fa hit miss
## 39.90 10.10 39.98 10.02
plotFit(fit) # graphical test
```

Observed (red) and predicted (boxplot) mean frequencies



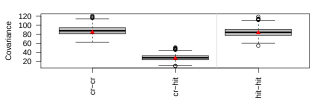
Model Fit: Predicted vs. Observed Data

■ Model fit for *covariance* of observed frequencies

```
cov(frequencies) # observed covariance matrix that is tested

## cr fa hit miss
## cr 84.29592 -84.29592 26.63061 -26.63061
## fa -84.29592 84.29592 -26.63061 26.63061
## hit 26.63061 -26.63061 83.32612 -83.32612
## miss -26.63061 26.63061 -83.32612 83.32612
plotFit(fit, stat = "cov") # graphical test
```

Observed (red) and predicted (gray) covariances



Model Fit: Predicted vs. Observed Data

Testing model fit

- The statistics T1 and T2 quantify the discrepancy between observed and expected means/covariances (similar to Pearson's X²)
- Posterior predictive p-values
 - Values around .50 indicate good model fit
 - Values close to 0 (or close to 1) indicate misfit
 - In contrast to frequentist p-values, PPP values are not uniformly distributed when generating data from the correct model

```
PPP(fit. M = 2000. nCPU = 4)
    ## ## Mean structure (T1):
       Observed = 0.03225169 : Predicted = 0.03227619 : p-value = 0.495
    ##
    ## ## Covariance structure (T2):
       Observed = 6.777547; Predicted = 6.895092; p-value = 0.504
    ##
    ## ## Individual fit (T1):
                                  5
                                        6 7 8
                                                              10
    ## 0.509 0.524 0.517 0.518 0.567 0.543 0.497 0.552 0.461 0.516 0.492 0.537
          13
               14
                     15
                           16
                                 17
                                       18
                                             19
                                                   20
                                                        21
                                                              22
                                                                          24
    ##
    ## 0.526 0.515 0.479 0.536 0.458 0.528 0.446 0.510 0.256 0.338 0.524 0.516
          25
               26
                     27
                           28
                                 29
                                       30
                                             31
                                                   32
                                                        33
                                                                          36
    ## 0.516 0.324 0.534 0.330 0.514 0.520 0.410 0.482 0.495 0.502 0.464 0.490
         37
                     39 40
                                 41
                                       42
                                            43
                                                   44
                                                        45
    ## 0.402 0.502 0.442 0.237 0.334 0.526 0.484 0.483 0.534 0.500 0.448 0.532
Daniel ##Heck 49
```

Summary

Modeling with TreeBUGS is simple

- Define model and clean data
- Draw MCMC samples
- Check convergence
- 4 Check model fit
- Interpret/plot parameters

Note that these are the usual steps in any Bayesian analysis. . .

Advanced Modeling

Within-Subject Comparisons

Assessing within-subject differences in parameters

- Data: Additional columns for separate within-subject conditions
- 2 Model: Write EQN file for within-subject design
- MCMC sampling (as usual)
- Comparison: Compute differences of parameters (transformed parameters)

(1) Data structure for within-subject design

```
freq_within <- read.csv("2htm_within.csv")
head(freq_within, 3)</pre>
```

```
high_cr high_fa high_hit high_miss low_cr low_fa low_hit low_miss
##
## 1
          44
                    6
                            36
                                        14
                                               41
                                                               40
                                                                         10
## 2
          38
                   12
                            39
                                        11
                                               45
                                                               38
                                                                         12
## 3
          43
                            45
                                               34
                                                      16
                                                               30
                                                                         20
```

Within-Subject Comparisons

(2) Model: Function for writing within-subject EQN files

- TreeBUGS has a function extends a standard MPT model to multiple within-subject conditions
- Essentially, model equations are copied and each parameter gets a new label (e.g., d_condition1)

```
##
                 Category
                                 Equation
            Tree
## 1
     high_target
                 high_hit
                                   d_high
## 2
     high_target high_hit (1-d_high)*g
     high_target high_miss (1-d_high)*(1-g)
## 3
## 4
       high_lure high_cr
                                   d_high
## 5
    high_lure high_fa (1-d_high)*g
       high lure high cr (1-d high)*(1-g)
## 6
## 7
      low target low hit
                                    d low
## 8
      low target low hit
                              (1-d low)*g
## 9
      low_target low_miss (1-d_low)*(1-g)
## 10
        low lure
                   low cr
                                    d low
## 11
      low lure low fa
                              (1-d low)*g
## 12
        low lure
                 low cr
                          (1-d low)*(1-g)
```

Within-Subject Comparisons

(4) Transformed parameters

- Often the interest is in the difference of a parameter across conditions

 Example: Difference in memory strength $\Delta_d = d_{\text{high}} d_{\text{low}}$
- Based on the MCMC samples, we can simply compute any function of interest
- We get a new set of posterior samples that can be summarized as usual
 # fit to all conditions:
 fit_within <- traitMPT("2htm_within.eqn", "2htm_within.csv")

 # compute difference in d:
 diff_d <- transformedParameters(
 fit_within,
 transformedParameters = list("diff_d = d_high d_low"),
 level = "group")
 summary(diff_d)\$statistics</pre>

```
## Mean SD Naive SE Time-series SE
## 0.3212361182 0.0345022709 0.0003319983 0.0010837807
```

Comparisons in between-subject designs

- Fit MPT model to each condition separately
 - \blacksquare Separate group-level parameters μ and Σ per group
- Compute differences in group-level parameters across conditions/models

```
## Mean SD 2.5% 50% 97.5% Time-series SE n.eff Rhat R_95% ## d.m1-d.m2 0.259 0.088 0.085 0.259 0.431 0.005 313 1.013 1.038 ## d.m1>d.m2 0.999 0.029 1.000 1.000 1.000 0.000 4967 1.227 1.264
```

Covariates: Correlations

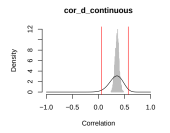
Correlation of MPT parameters with external covariates

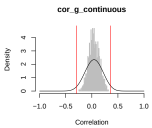
- New argument covData: A data frame or file name with values of the covariate(s)
- TreeBUGS computes the correlation of these covariates with the latent person parameters θ (probit values)

Covariates: Correlations

- Note that the posterior samples of the (descriptive) correlations only reflect uncertainty with respect to the MPT parameters
- We also need to consider the number of participants (sample size)!
- Solution: Use an analytical solution or the posterior distribution of the correlation (Ly et al., 2018)

correlationPosterior(fit cor)





```
## 2.5% 50% 97.5%
## cor_d_continuous 0.06 0.335 0.57
## cor g continuous -0.29 0.035 0.36
```

Regression of MPT parameters on covariates

- lacktriangle Example: Predict memory performance d as a function of age
- Statistically, this requires an regression extension to the model
- The latent probit values θ'_i are predicted by a design matrix X:

$$\theta_i' = \mu + X_i \beta + \delta_i$$

Implementation in TreeBUGS

Requires only two new arguments to provide the data (age of persons) and the regression structure (predict parameter D_n by age)

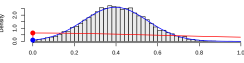
```
## Mean SD 2.5% 50% 97.5% n.eff Rhat R_95% ## 0.18 0.07 0.04 0.18 0.31 301.00 1.02 1.06
```

Bayes Factor for Covariate

Compute a Bayes factor

- H0: Slope parameter $\beta = 0$
- H1: Slope parameter $\beta \sim \text{Cauchy}(0, r)$ (with scale parameter r)
- Method: Savage-Dickey density ratio (Wagenmakers, 2010)
 - Bayes factor H1 vs. H0: prior devided by posterior density (at $\beta = 0$)
 - Only works for simple regression with 1 predictor (Heck 2019)

Bayes factor B_10=6.787 (prior red; posterior blue)



Standardized slope parameter: d ~ continuous

```
## BF_0> BF_>0
## slope_d_continuous 0.1473361 6.787201
```

Between-subject designs: Assumptions about the covariance matrix Σ

- lacktriangle Separate covariance matrix per condition: $\Sigma_1, \Sigma_2, \dots$
 - See previous slides: betweenSubjectMPT(fit1, fit2)
- B) Identical covariance matrix Σ across conditions
 - Similar to ANOVA: "pooled variance" (Rouder & Morey; 2012)
 - \blacksquare Manipulation only affects the mean parameters μ

```
## Mean SD 2.5% 50% 97.5% p(one-sided vs. overall)
## d_discrete[group_a] 0.63 0.09 0.46 0.63 0.79 0.47
## d_discrete[group_b] 0.65 0.08 0.48 0.65 0.80 0.47
```

Sensitivity/robustness analysis

Define different priors

- Prior distributions in the latent-trait MPT necessary for:
 - Latent (probit-) mean μ
 - Latent (probit-) covariance matrix Σ : scaled inverse Wishart with
 - Prior matrix V
 - Degrees of freedom df
 - \blacksquare Scaling parameter ξ
- Example: We assume that guessing probabilities are around 50%

Understanding Priors

What do the priors actually mean?

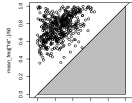
- Draw samples from the prior
- Plot mean/SD of MPT parameters

Prior on group mean: mu=dnorm(0,1) Prior on group mean: mu=dnorm(0,5) 0.8 1.5 1.0 0.4 0.5 0.4 0.6 0.0 0.2 0.4 0.6 0.8 Group mean (probability scale) Group mean (probability scale) Prior on group SD: xi=dunif(0.2) Prior on group SD: xi=dunif(0.2) 0.4 0.4 0.2 0.2 0.0

Prior Predictive Sampling

Prior predictive distribution

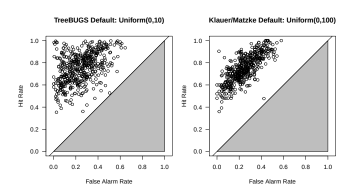
- Draw samples from the prior
- Draw new data (response frequencies)
- 3 Assess predicted data (e.g., plots or descriptive statistics)



Prior Predictive Sampling

Excursion: Different default priors for the scale parameter ξ

- **I** TreeBUGS (Heck et al., 2018): $\xi \sim \text{Uniform}(0, 10)$
- **2** Klauer (2010) and Matzke et al. (2015): $\xi \sim \text{Uniform}(0, 100)$



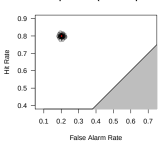
Posterior Predictive

Posterior Predictive Distribution

- What data does the fitted model predict?
- Use posterior samples of the parameters to draw new samples of the data (i.e., predicted response frequencies)
 - Note: These are the basis of posterior-predictive checks (T1 and T2 statistics)

```
postpred <- posteriorPredictive(fit, M = 100, nCPU = 4)</pre>
```

Posterior predicted (mean frequenci



Sensitivity and robustness analysis

- Assessing the impact of priors, estimate necessary sample size for specific analysis etc.
 - I Generate data from (correct or wrong) model
 - 2 Fit model
 - 3 If necessary: Replicate with a for-loop

Appendix

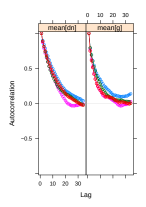
Appendix: Testing for Heterogeneity

Appendix: Convergence

Autocorrelation function

- \blacksquare How strongly are the MCMC samples correlated between iteration t and iteration t+Lag?
- Ideally, these curves should rapidly decrease towards zero.

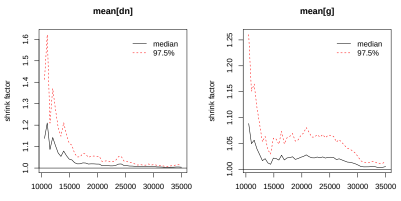
```
plot(fit, parameter = "mean", type = "acf")
```



Appendix: Convergence

- Plot evolution of Gelman-Rubin statistic
 - Also known as: "potential scale reduction factor" or "R hat"
 - Similar to ANOVA: Compares between-chain and within-chain variances (large differences between these variances indicate nonconvergence)
 - Statistic should be close to 1

```
plot(fit, parameter = "mean", type = "gelman")
```



Daniel W. Heck last iteration in chain last iteration in chain 4

Appendix: MPT Versions Implemented in TreeBUGS

MCMC samplers available in TreeBUGS

- Fixed-effects ("standard") MPT: Gibbs sampler in C++
- Beta-MPT: JAGS and C++
- Latent-trait with extensions: JAGS
 - Combination of random- and fixed-effects parameters
 - Continuous and categorical covariates
 - Group-level structure: Independent normal distributions

References I

Heck, Daniel W. 2019. "A Caveat on the Savage-Dickey Density Ratio: The Case of Computing Bayes Factors for Regression Parameters." *British Journal of Mathematical and Statistical Psychology* 72: 316–33. https://doi.org/10.1111/bmsp.12150.

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Klauer, K. C. 2010. "Hierarchical Multinomial Processing Tree Models: A Latent-Trait Approach." *Psychometrika* 75: 70–98. https://doi.org/10.1007/s11336-009-9141-0.

Matzke, Dora, Conor V. Dolan, William H. Batchelder, and Eric-Jan Wagenmakers. 2015. "Bayesian Estimation of Multinomial Processing Tree Models with Heterogeneity in Participants and Items." *Psychometrika* 80: 205–35. https://doi.org/10.1007/s11336-013-9374-9.

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